

# Genetic Variation in Odorant Receptors Contributes to Variation in Olfactory Behavior in a Natural Population of *Drosophila melanogaster*

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Genbank accession numbers for *D. melanogaster* sequences: *Or22a/Or22b*: JF745070–JF745092, JF719972–JF719994, JF719957–JF719971; *Or35a*: JF719919–JF719956; and *Or47a*: JF745032–JF745069.

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## Abstract

Chemoreception is a principle modality by which organisms gain information from their environment, and extensive variation in odor-mediated behavior has been documented within and among species. To examine the mechanisms by which sensory systems mediate these responses, we ask to what extent variation in *Drosophila melanogaster* odorant receptor genes contributes to variation in odor-mediated behavior. Significant differences in behavioral responses to structurally similar odorants, methyl hexanoate and ethyl hexanoate, were found in a natural population. Polymorphisms in 3 genomic regions (*Or22a/Or22b*, *Or35a*, and *Or47a*) were identified and associated with variation in behavior to these esters. Overall similarity in association profiles for both odorants was observed, except for *Or47a* in which polymorphisms were associated solely with variation in responses to ethyl hexanoate. Our analyses were then extended to examine polymorphisms in 3 odorant receptors previously reported to contribute to variation in olfactory behavior for the chemically distinct odorants benzaldehyde and acetophenone. Two *Or10a* polymorphisms were associated with variation in response to ethyl hexanoate. Finally, differences in *Or35a* and *Or47a* expression were associated with variation in responses to ethyl hexanoate. These results demonstrate that the genetic variation at the peripheral sensory stage plays a role in mediating differences in odor-mediated behavior.

**Key words:** behavior, chemosensory, olfaction, single nucleotide polymorphism

## Introduction

Understanding the mechanisms by which sensory systems mediate behavioral responses to stimuli is a fundamental goal in behavioral neuroscience, and numerous advances have been made uncovering both genetic mechanisms and neural circuitry involved in sensory transduction and perception. In olfaction, for instance, it is well established that chemicals are bound by a diverse family of receptor proteins expressed in olfactory receptor neurons (ORNs), whose combinatorial activation results in a spatial pattern of brain activity (Buck and Axel 1991; Vassar et al. 1994; Mombaerts et al. 1996; Laissue et al. 1999; Gao et al. 2000; Vosshall et al. 2000; Bhalarao et al. 2003; Wang et al. 2003). This neural code enables an organism to identify and discriminate among chemical signals in its environment. However, despite these advances, it remains unclear how underlying molecular polymorphisms in these receptors contribute to variation in olfaction-mediated behavior within and between species and the nature and pattern of polymorphisms contributing to receptor–ligand specificity.

The *Drosophila melanogaster* olfactory system affords us the opportunity to begin to address this issue because odorant receptor loci have been identified and many of their response spectra have been characterized (Clyne et al. 1999; Gao and Chess 1999; Vosshall et al. 1999; Dobritsa et al. 2003; Hallem et al. 2004; Hallem and Carlson 2006). In *Drosophila*, ORNs are located within sensilla distributed over the third antennal segments and maxillary palps. Each sensillum contains up to 4 neurons with specific combinations of neurons restricted to distinct spatial antennal regions (de Bruyne et al. 1999, 2001). Each ORN typically expresses a single odorant receptor, and thus, the response profile of individual ORNs is determined by the individual odorant receptor expressed, with the majority of receptors responding to numerous and overlapping suites of ligands (Dobritsa et al. 2003; Wang et al. 2003; Hallem et al. 2004; Hallem and Carlson 2006).

The *D. melanogaster* odorant receptor (*Or*) family consists of 62 seven transmembrane proteins transcribed from 60 *Or*

genes (Clyne et al. 1999; Vosshall et al. 2000; Robertson et al. 2003; Vosshall and Stocker 2007). The membrane topology of these receptors is opposite that of vertebrate odorant receptors, with the N-terminus in the cytoplasm of the ORN (Benton et al. 2006). Evidence also suggests that these odorant receptors dimerize with a highly conserved noncanonical receptor, *Orco* (formerly known as *Or83b*), that serves to localize odorant receptors in the sensory dendritic membrane (Jones et al. 2005; Benton et al. 2006) and form a class of ligand-gated nonselective cation channels (Sato et al. 2008; Wicher et al. 2008). Most recently, studies have also identified odorant receptor function in a group of ionotropic glutamate receptor-related proteins expressed in a subset of ORNs devoid of other previously identified odorant receptors (Benton et al. 2009; Abuin et al. 2011).

Alterations in both the central and the peripheral nervous systems contribute to extensive variation in olfaction-mediated behavioral responses within and among *Drosophila* species (Fuyama 1976, 1978; Alcorta and Rubio 1988, 1989; Mackay et al. 1996; Wang et al. 2007; Rollmann et al. 2010; Wang et al. 2010). In a comparative study between *D. melanogaster* and *D. simulans*, for example, differences in the specificity of particular ORNs was observed, with a shift in ligand affinity from ethyl hexanoate to methyl hexanoate (Stensmyr et al. 2003). Since the response profile of individual ORNs is determined by the odorant receptor expressed, Stensmyr et al. (2003) suggested that this shift may be due to a substitution in the ligand-binding domain of the receptor. Differences in sensitivity to these 2 odorants were also observed between *D. melanogaster* and *D. sechellia* with changes in sensitivity thought to be due to coding sequence changes in receptors as well as overexpression of methyl hexanoate-responsive neurons (Stensmyr et al. 2003; Dekker et al. 2006). Within species, polymorphisms in odorant receptor loci have also been implicated in mediating variation in olfactory behavior (Rollmann et al. 2010). In a natural *D. melanogaster* population, sequence variation in 3 odorant receptor loci, *Or10a*, *Or43a*, and *Or67b*, was associated with variation in behavioral responses to structurally similar odorants, benzaldehyde and acetophenone. Association profiles for both odorants revealed that distinct polymorphic sites in these loci contribute to differences in behavior and that some but not all of the same polymorphisms were associated with behavioral responses to both odorants (Rollmann et al. 2010).

Here, we examine how molecular polymorphisms in odorant receptor loci contribute to variation in behavior within a natural *D. melanogaster* population. We first assess the extent of variation in attraction/avoidance to 2 esters, ethyl hexanoate and methyl hexanoate, which are similar in structure, differing only by a single carbon length. These esters have been shown to be physiologically active and present in fruit extracts, the natural feeding/breeding substrate of *Drosophila* (Stensmyr, Giordano, et al. 2003). A significant difference in olfactory sensitivity to these esters has been previously observed among *Drosophila* species (Stensmyr et al. 2003; Dekker et al. 2006), but within-species variation had

yet to be examined. Next, we identify numerous single nucleotide polymorphisms (SNPs) and insertion and deletion polymorphisms (indels) in 3 *D. melanogaster* odorant receptor loci, *Or22a*, *Or35a*, and *Or47a* that have been shown to be electrophysiologically tuned to these esters (Dobritsa et al. 2003; Hallem et al. 2004; Hallem and Carlson 2006). We also sequence *Or22b* due to its close physical proximity to *Or22a* (540 bp downstream) on the second chromosome, its coexpression in the same cell, and the previously reported presence of a single chimeric *Or22ab* gene within some *D. melanogaster* populations (Dobritsa et al. 2003; Aguadé 2009). Subsequently, we conduct genotype–phenotype association analyses to determine which polymorphic sites (if any) in these 4 *Or* loci contribute to variation in olfactory behavior and examine whether lines with overall more positive versus negative behavioral response indices to ethyl hexanoate show differences in *Or* gene expression levels.

Finally, we extend our analyses of genotype–phenotype relationships to assess whether genetic variation in 3 previously published odorant receptor loci, *Or10a*, *Or43a*, and *Or67b* (Rollmann et al. 2010) also contributes to the observed behavioral differences. Polymorphisms in these loci have been previously implicated in mediating variation in behavioral responses to odorants, benzaldehyde and acetophenone (Rollmann et al. 2010). We predict that no genotype–phenotype associations will be observed with variation in *Or43a* and *Or67b* because no ligand specificity to ethyl hexanoate or methyl hexanoate has been documented. However, ethyl hexanoate has been previously demonstrated to elicit inhibitory responses in *Or10a* expressing neurons. Our results demonstrate that changes at the most peripheral sensory stage contribute to within-species differences in behavioral responses to environmental stimuli.

## Materials and methods

### Experimental animals

Isofemale lines were derived from a natural population of *D. melanogaster* collected in Raleigh, NC. Each line was subsequently inbred by 20 generations of full-sib mating (Ayroles et al. 2009). Flies were raised on a cornmeal/agar/molasses media at 25 °C with a 12 h light–dark cycle and behaviorally tested 3- to 7-day posteclosion.

### Chemicals

Ethyl hexanoate and methyl hexanoate were purchased from Sigma-Aldrich at the highest purity available (>99% purity). Odorants were diluted in paraffin oil (Sigma-Aldrich).

### T-maze olfactory assay

Behavioral assays were essentially in accordance with Helfand and Carlson (1989) with minor modification. Thirty flies were placed into the T-maze apparatus and allowed to

acclimate to conditions in a holding chamber for 1 min. Flies were then given a choice between airflow (500 mL/min) from the 2 arms of the maze, one containing the diluted odorant and one solely containing the paraffin oil vehicle. Flies were allowed 1 min to choose between the odorant and the control side. The response index (RI) was calculated by the formula:  $RI = (\text{No. flies choosing odor} - \text{No. flies choosing control}) / (\text{No. total flies})$ . A positive RI represents attraction to the odor (with a maximal attractive response of +1), whereas a negative RI indicates repulsion (with a maximal repulsion response of -1). To optimally resolve variation among wild-derived lines, dose response curves were first generated for each odorant on a subset of 4 randomly chosen wild-derived lines. Subsequent behavioral assays on the remainder of the wild-derived lines were then performed at 0.1% (v/v) methyl hexanoate and 1.0% (v/v) ethyl hexanoate. Approximately 9 replicate tests were performed per line and sex. *Drosophila* lines were tested using a randomized design, and all assays were conducted in the morning.

### Statistical analyses of behavior

Data were analyzed using JMP 8.0 (SAS Institute). Comparisons among lines were made using a 2-way factorial analysis of variance (ANOVA) model  $y = \mu + L + S + L \times S + E$ , where sex ( $S$ ) is the fixed effect of sex, line ( $L$ ) and  $L \times S$  are the random effect of line and line by sex, respectively, and  $E$  indicates error. To examine if there was a correlation between the behavioral responses to ethyl hexanoate and methyl hexanoate, the mean response indices for all lines and sexes were plotted. The response index for methyl hexanoate was arbitrarily chosen as the independent variable and the response index for ethyl hexanoate was chosen as the dependent variable.

### Identification of polymorphisms in odorant receptor loci

Genomic DNA was extracted using the Puregene DNA purification system (Qiagen) for the same wild-derived lines for which behavioral data were collected. Primers were designed for 4 odorant receptors (*Or22a*, *Or22b*, *Or35a*, and *Or47a*) in order to obtain overlapping fragments encompassing the full-length coding and noncoding regions and ~700 bp of the 5' and 3' untranslated regions. Polymerase chain reaction (PCR) was performed, and products sequenced with original and nested PCR primers. Sequences were analyzed using Vector NTI Advance 11 software (Informax). DNA sequences for *Or10a*, *Or43a*, and *Or67b* were obtained previously (Rollmann et al. 2010; *Or10a*,  $N = 79$ ; *Or43a*,  $N = 72$ ; and *Or67b*,  $N = 60$ ).

### Genotype–phenotype association analyses

Associations between individual polymorphic sites in each odorant receptor locus and variation in behavioral responses to ethyl and methyl hexanoate were determined by 2-way factorial ANOVA, model  $y = \mu + M + S + M \times S + E$ , in which  $\mu$  is the overall mean, the molecular marker ( $M$ )

and sex ( $S$ ) are fixed effects, and  $E$  represents error. A conservative Bonferroni correction was performed to account for multiple tests. Likewise, when multiple polymorphisms in a given receptor were associated with variation in behavior, haplotypes were determined, and associations between haplotype and variation in behavior was assessed according to the 2-way factorial model  $y = \mu + H + S + H \times S + E$ , in which  $\mu$  is the overall mean, the haplotype ( $H$ ) and sex ( $S$ ) are fixed effects, and  $E$  represents error. Polymorphisms present in only one line were not included in the association analyses. Linkage disequilibrium (LD) between markers was determined using Tassel 2.1 software ([www.maizegenetics.net/tassel](http://www.maizegenetics.net/tassel)) and examined using Fisher's Exact test. An  $R^2$  value of 1 was indicative of complete LD, and polymorphisms in complete LD were treated as a single linkage group in Bonferroni calculations. Thirty-eight wild-derived lines were examined for each odorant receptor locus.

### Genotyping

To assess support for the significant genotype–phenotype relationships in a larger sample, SNPs or indels identified as significantly associated with natural variation in olfactory behavior in our analyses of 38 wild-derived lines were genotyped in 82 additional lines from the same population. The genotyping approach was dictated by the number of polymorphisms significantly associated with behavior for a given locus and the pattern of LD. Either DNA sequencing of the genomic region of interest or Taqman Sample-to-SNP kit based methods (Applied Biosystems) was used. The presence or absence of the *Or22ab* chimera was determined by independent design of 2 primer pairs: One pair had one of the primers located within the deleted region (*Or22a*-F: GGTGTGAAGGTGTAGTTTGTAG, *Or22a*-R: TATCAAGCGGTGATCTCG). The second pair had primers spanning the deleted region (*Or22a*-F2: GACGCTTCACTCTCCGTG, *Or22a*-R2: GATTACTACTCGTAA-CAAGC). Taqman primers and probes were as follows: *Or35a*-F: GGGAGCAAATCCTGCAGTACT, *Or35a*-R: GGCCAAGTTAATGAGCTTTAGTAATCG, *Or35a*-variant1: VIC-TCGCTGGGATTTGTAG, *Or35a*-variant2: FAM-CGCTGGGACTTGTAG, *Or47a*-F: CGCACGGCGATGTCCTA, *Or47a*-R: GCA-CAAAAATCAATCCAAAGCCTTGT, *Or47a*-variant1: VIC-ATGCTGAGATCATTCTCCT, and *Or47a*-variant2: FAM-TGCTGAGATCCTTCTCCT.

### Gene expression

Total RNA was isolated from 50 *D. melanogaster* heads using Trizol (Invitrogen) and DNase treated (Ambion) according to manufacturers' instructions. Total RNA for 3 independent samples per sex and line was isolated. First strand cDNA was then synthesized using the AccuScript High Fidelity First Strand cDNA synthesis kit (Agilent Technologies). Quantitative real-time PCR assays were conducted

with an ABI 7300 and using Taqman chemistry (Applied Biosystems). For each independent RNA isolation for each line and sex, 2 technical replicate measurements were made. Taqman gene expression assays for each gene were as follows: *Or22a*, Dm01814175\_g1; *Or22b*, Dm01844971\_g1; *Or35a*, Dm01807569\_g1; *Or47a*, Dm01818125\_g1; and *Rpl32* (endogenous control), Dm02151827\_g1. The  $\Delta\Delta C_t$  method was used for comparison of relative gene expression.

## Results

### Behavioral responses to select esters and behavioral correlations

Behavioral variation was quantified among the wild-derived inbred lines to assess the extent of naturally occurring variation in olfactory behavior to 2 esters, ethyl hexanoate and methyl hexanoate. Significant phenotypic variation was detected among the wild-derived lines in response to both odorants (MetHex: degrees of freedom [df] = 37,  $F = 6.16$ ,  $P < 0.0001$ ; EtHex: df = 119,  $F = 6.24$ ,  $P < 0.0001$ ; Figure 1A and B, respectively). The behavioral response indices were broad and ranged from 0.394 to  $-0.589$  for methyl hexanoate and 0.278 to  $-0.702$  for ethyl hexanoate (Figure 1A and B). No significant difference was observed between the sexes for behavioral response to either odorant (MetHex: df = 1,  $F = 0.31$ ,  $P = 0.58$ ; EtHex: df = 1,  $F = 0.66$ ,  $P = 0.42$ ).

There was a significant positive correlation between responses to ethyl and methyl hexanoate ( $F = 69.13$ ,  $P < 0.0001$ ,  $R = 0.695$ ; Figure 2). This was expected because methyl hexanoate and ethyl hexanoate have similar chemical structures (differing solely in their carbon length), and electrophysiological response profiles for several odorant receptor loci often revealed overlapping sensitivity to both odorants (Dobritsa et al. 2003; Hallem et al. 2004; Hallem and Carlson 2006). Our results demonstrated that significant intraspecific variation in olfactory behavior exists in this natural (Raleigh, NC) population and that odorants of similar chemical structure elicit comparable behavioral responses among lines.

### Identification of molecular polymorphisms

Four receptors (*Or22a*, *Or22b*, *Or35a*, and *Or47a*) were sequenced for 38 inbred lines, derived from a natural population, for which behavioral responses to ethyl hexanoate and methyl hexanoate were measured. Three of these receptors (*Or22a*, *Or35a*, and *Or47a*) have been previously shown to respond strongly to esters (Hallem et al. 2004; Hallem and Carlson 2006), and *Or22b* was sequenced as it is physically located close to *Or22a*.

Multiple SNPs and indels were identified in each of the 4 odorant receptors (Supplementary Table 1 and Figure 3). The  $\sim 2563$ -bp region sequenced for *Or35a* had a total of 50 polymorphisms, 4 of which result in amino acid substitutions (Lys38Ile, Leu47Val, Thr250Ile, and Asn348Ser). The

$\sim 2549$ -bp sequence for *Or47a* had a total of 164 polymorphisms, with one resulting in an amino acid substitution (Val1176Ile). Because the *Or22a* and *Or22b* regions are in close physical proximity they were considered together. This  $\sim 4460$  bp sequenced gene region contained 68 polymorphisms, which results in 11 amino acid substitutions in *Or22a* (His8Tyr, Val30Leu, Glu38Val, Trp51Leu, Met59Ile, Ile67Met, Met93Val, Cys100Ser, Glu196Ala, Asp201Glu, and Tyr382Ala) and 6 in *Or22b* (Ile25Val, Asn92Asp, Arg194Met, Asp201Ala, Val213Leu, and Leu354Ile). One polymorphism, a  $\sim 2090$ -bp deletion encompassing the last 3 exons of *Or22a* and the first exon of *Or22b*, resulted in a chimeric *Or22ab* gene (Figure 3B) with a frequency of 0.4 in the population sample. This *Or22ab* chimera has been reported previously by Aguadé (2009) and has been shown to respond strongly to esters (de Bruyne et al. 2010).

### Linkage disequilibrium

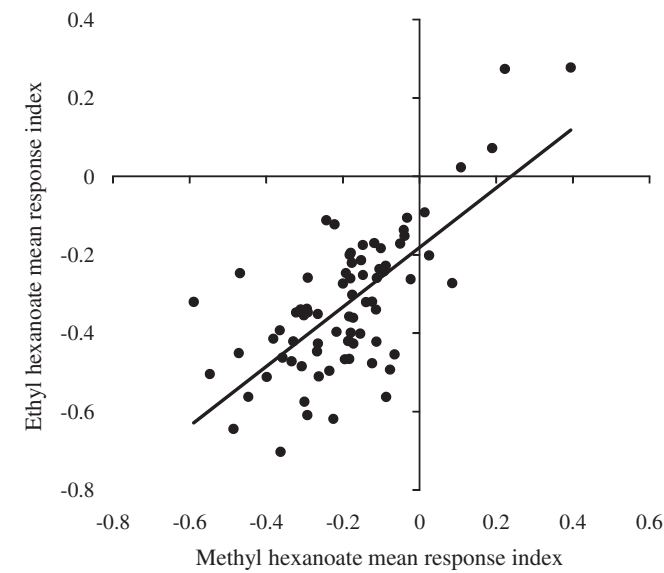
LD analyses were performed for the 4 odorant receptor loci to detect nonrandom associations among polymorphic sites (Figure 4). As in previous analyses, LD analyses of the *Or22a* and *Or22b* gene regions were conducted together. For each odorant receptor gene region, LD generally decayed rapidly with physical distance in a manner consistent with mutation-drift equilibrium so that significant LD was generally observed only between polymorphisms in close physical proximity (Figure 4). However, 3 incidences of long-range LD were observed in the *Or22a/Or22b* complex. LD spanned a  $\sim 3$ -kb region with C4T and C22T in *Or22a* and T860A, 909 Del (9 bp), and T960C in *Or22b* in complete LD. Long-range LD (across  $\sim 2.4$  kb) was also observed between A27T, C60G, and C61A in *Or22a* and C181G, G183A, A184C, T185C, G282A, and A334G in *Or22b* and between a third group consisting of A113T, A140G, C143C in *Or22a* and C955A in *Or22b*, spanning a  $\sim 2.9$ -kb region.

### Genotype–phenotype associations

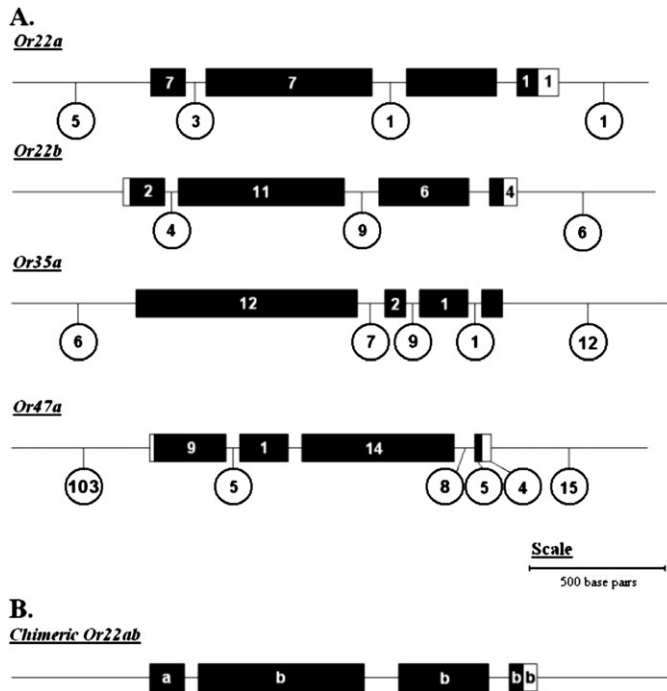
To determine the extent and nature of molecular polymorphisms in these receptors which may contribute to variation in olfaction-mediated behavior, we analyzed associations between sequence variation in 3 odorant receptor gene regions (*Or22a/Or22b*, *Or35a*, and *Or47a*) and variation in behavioral responses to 2 odorants, ethyl hexanoate and methyl hexanoate (Figure 5). In the *Or22a/Or22b* gene region, we observed 2 polymorphisms that exceeded a conservative Bonferroni correction for multiple tests and were associated with variation in responses to both methyl hexanoate and ethyl hexanoate (Figure 5A). One of these was a silent substitution in the second exon of *Or22b* (T240C) and the other was a 16-bp insertion (1541 Ins) in the 3' untranslated region of *Or22b*, a region highly conserved among closely related *Drosophila* species (*D. simulans*, *D. sechellia*, and *D. yakuba*; *Drosophila* 12 Genomes Consortium 2007, <http://genome>.







**Figure 2** Correlation between behavioral responses to ethyl hexanoate and methyl hexanoate. The mean response indices for ethyl hexanoate for each sex and line are plotted against the behavioral response indices for methyl hexanoate.



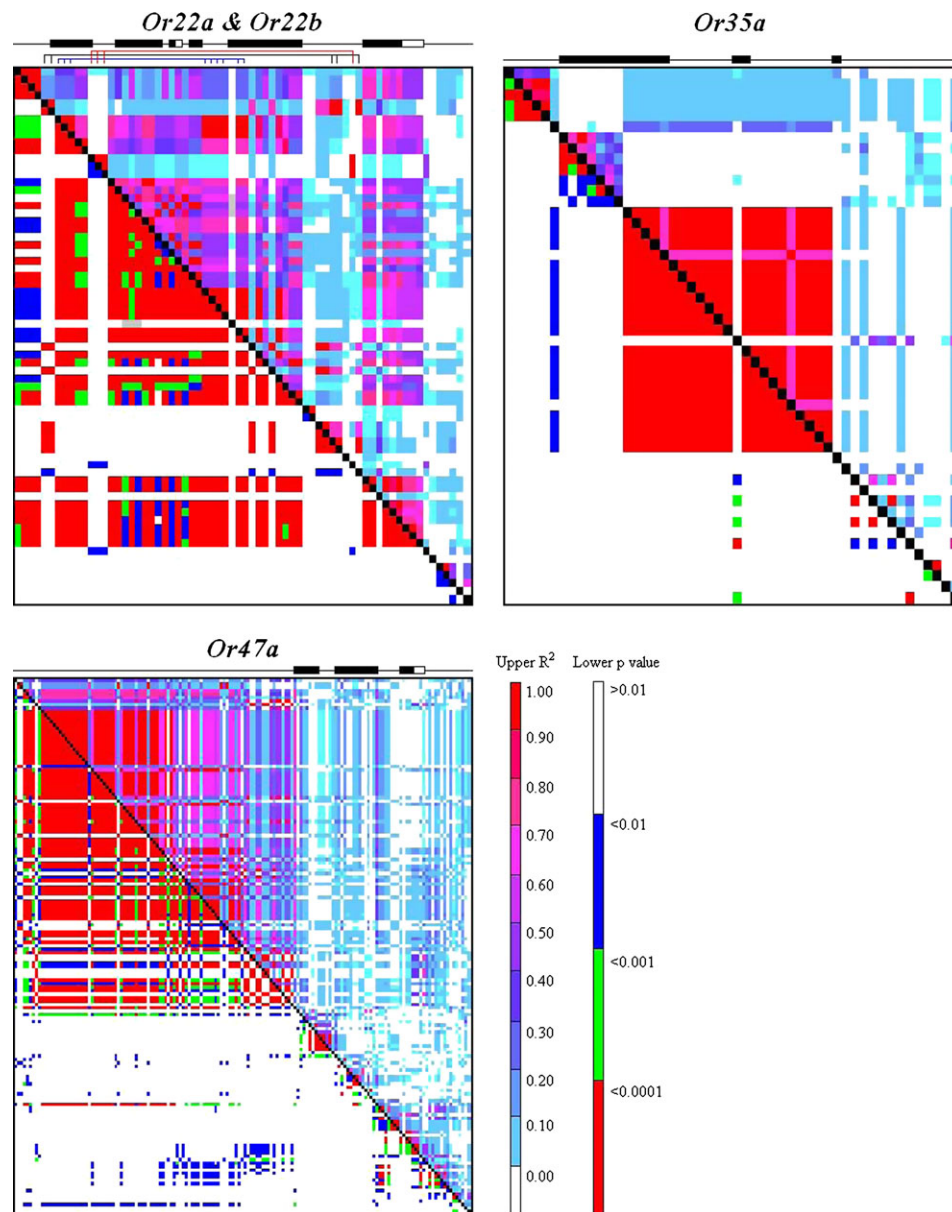
**Figure 3** A schematic of the odorant receptor gene structure and molecular polymorphisms in each *Or* gene region. (A) The genomic region is depicted with a solid line with black boxes representing exons and white boxes untranslated regions. The number of polymorphic markers in each region is depicted numerically either directly within the exons or in circles underneath. (B) For the chimeric *Or22ab* gene, the source of each exon is indicated by an “a” for *Or22a* or a “b” for *Or22b*.

ucsc.edu/cgi-bin/hgGateway). In the case of *Or35a*, 2 polymorphisms (G-455A and A1195G) were significantly associated with behavioral responses to both odorants (Figure 5B).

The G-455A polymorphism is located directly upstream of the initiation codon for *Or35a* between 2 highly conserved blocks of sequence. The A1195G is located in the third exon, which results in an amino acid substitution (Asn348Ser) near the C-terminus of the receptor. Finally, 7 polymorphic markers (G195T, T1300C, C1321T, A1324G, A1336C, 1349Del2bp, and C1357T) in *Or47a* were significantly associated with behavioral responses only to ethyl hexanoate (Figure 5C). G195T is a silent substitution located in the first exon of *Or47a*. T1300C, C1321T, A1324G, and A1336C are silent substitutions located in exon 4, with T1300C, C1321T, and A1324G in complete LD (Figure 4). An A1336C polymorphism and 2 markers (1349Del2bp and C1357T) in the untranslated region of exon 4 are in complete LD (Figure 4). The polymorphisms (1349Del2bp and C1357T) are in close proximity to the termination codon ( $\leq 12$  bp away), and comparisons among closely related species (*D. simulans*, *D. sechellia*, *D. yakuba*, and *D. erecta*) indicate that they are located within a conserved region. No polymorphisms in *Or47a* were significantly associated with variation in behavioral responses to methyl hexanoate.

Because each odorant receptor contained multiple polymorphisms associated with variation in olfactory behavior, we also conducted haplotype analyses to determine the potential combined effects of polymorphisms on behavior. For polymorphisms associated with behavioral responses to methyl hexanoate, 3 haplotypes were identified for the *Or22a/Or22b* region and 3 for *Or35a* (Table 1 and Figure 6A). For the *Or22a/Or22b* region haplotypes, a significant avoidance of methyl hexanoate was observed for haplotype 3 relative to haplotypes 1 and 2 ( $df = 2$ ,  $F = 21.14$ ,  $P < 0.0001$ ). Haplotype analysis of *Or35a* also revealed significant difference among haplotypes ( $df = 2$ ,  $F = 8.99$ ,  $P = 0.0003$ ). Similarly, for sequence variants associated with variation in behavioral responses to ethyl hexanoate, significant differences in mean responses among haplotypes were observed for *Or35a* ( $df = 2$ ,  $F = 14.27$ ,  $P < 0.0001$ ) and *Or47a* ( $df = 4$ ,  $F = 11.11$ ,  $P < 0.0001$ ), with 3 haplotypes identified for *Or35a* (the same as those for methyl hexanoate) and 5 for *Or47a* (Table 1 and Figure 6B). Of particular note is the significantly greater mean behavioral response of haplotype 4 relative to all other haplotypes.

We then extended the sample to 120 wild-derived lines for which measurements of behavioral responses to ethyl hexanoate and genotypes for focal polymorphisms were obtained. With the exception of the *Or47a* A1336C linkage group, the statistical association between the phenotypic means for behavioral responses to ethyl hexanoate and each examined focal polymorphic marker in the sample of 120 alleles continues to support the observed significant genotype–phenotype relationships for *Or35a* markers G-455A ( $df = 1$ ,  $F = 9.72$ ,  $P < 0.002$ ; frequency A allele, 0.13) and A1195G ( $df = 1$ ,  $F = 25.09$ ,  $P < 0.0001$ ; frequency of G allele, 0.05) and variation in behavioral responses to ethyl hexanoate. Significant associations were also observed for the T1300C

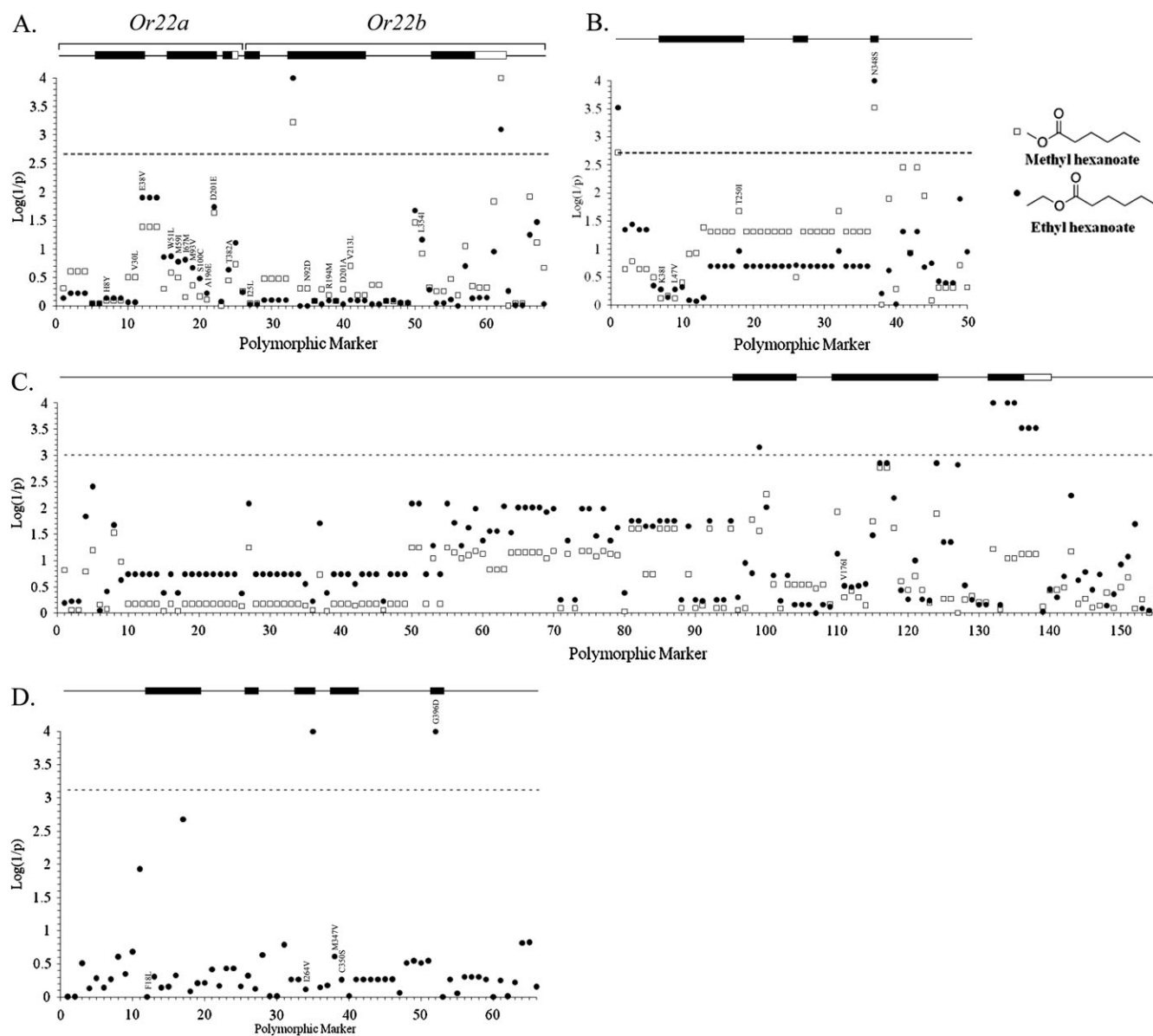


**Figure 4** LD plots between markers for each odorant receptor. The upper diagonal represents the  $R^2$  value, whereas the lower diagonal represents the  $P$  value for all marker combinations.  $R^2$  values of 1 indicate complete LD. The location of coding and noncoding regions for each gene is shown above each LD plot with coding regions indicated by black boxes and untranslated regions by white boxes. In the case of the *Or22a/Or22b* gene region, colored lines below the gene region highlight distinct regions of long-range LD.

marker in *Or47a* ( $df = 1$ ,  $F = 6.11$ ,  $P < 0.01$ ; frequency C allele, 0.15) and the T240C marker in the *Or22a/Or22b* region ( $df = 1$ ,  $F = 26.88$ ,  $P < 0.0001$ ; frequency C allele, 0.05). Finally, in regard to the deletion resulting in a chimeric *Or22ab* gene, we found the presence/absence of this  $\sim 2$ -kb deletion to be significantly associated with behavioral responses to ethyl hexanoate ( $df = 1$ ,  $F = 4.62$ ,  $P < 0.03$ , frequency 0.39).

Finally, we took advantage of previously published sequence data for *Or10a*, *Or43a*, and *Or67b* in these same wild-derived lines (Rollmann et al. 2010) to ask whether polymorphisms in these receptors were associated with variation in

behavioral responses to ethyl hexanoate. Consistent with the lack of *Or43a* and *Or67b* specificity to ethyl hexanoate, no associations were found between polymorphisms in these loci and variation in responses to ethyl hexanoate. We did, however, find genotype–phenotype associations between sequence variation in *Or10a*, a receptor for which inhibitory electrophysiological responses to ethyl hexanoate have been observed (Hallem and Carlson 2006) and variation in behavioral responses to ethyl hexanoate (Figure 5D). More specifically, the C1131T and G1481A polymorphisms contributed to behavioral variation. These polymorphisms are distinct from those associated with variation in behavioral



**Figure 5** Associations between each polymorphic marker and mean behavioral response index for (A) *Or22a* and *Or22b*, (B) *Or35a*, (C) *Or47a*, and (D) *Or10a*. Association between markers and olfactory responses to ethyl hexanoate and methyl hexanoate are represented by black circles or open boxes, respectively. Markers above the Bonferroni threshold (dashed black line) are significantly associated with behavioral variation. Above each graph is a schematic of the genomic region for each *Or* with coding regions indicated by a black box, nontranslated regions by a white box, and noncoding DNA segments by a line.

responses to benzaldehyde and acetophenone (Rollmann et al. 2010). The G1481A polymorphism results in a glycine to aspartic acid amino acid substitution (G396D).

### Gene expression

Molecular polymorphisms in noncoding regions can result in alteration to gene expression levels. To assess whether noncoding polymorphisms associated with variation in olfactory behavior were associated with gene expression differences in the *Or22a/Or22b* gene region, *Or35a*, and *Or47a*, we conducted quantitative real-time PCR experiments on lines

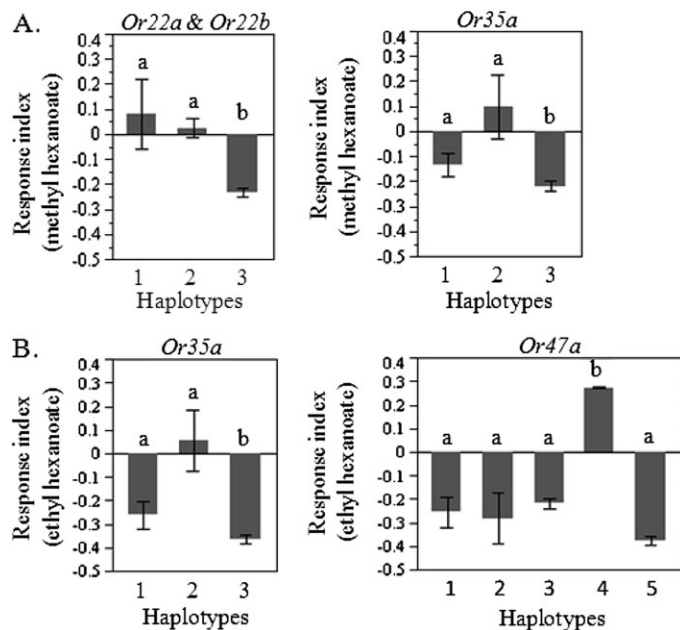
with overall more positive versus negative behavioral response indices to ethyl hexanoate. For each gene, 4 lines with more negative behavioral response indices to ethyl hexanoate and carrying a given allele and 4 lines with more positive indices carrying the alternate allele were selected. In the case of the *Or22a/Or22b* region, we did not observe significant differences in *Or22b* expression among lines with positive versus negative behavioral responses to ethyl hexanoate but confirmed the presence of the *Or22ab* chimera. However, for *Or35a* and *Or47a*, we did observe significant differences in gene expression levels among lines with alternate alleles and respective positive/negative behavioral response indices



**Table 1** Haplotype analyses for polymorphisms significantly associated with behavioral responses to methyl hexanoate or ethyl hexanoate for each locus

Gene region	Odorant	Number of haplotypes	Haplotype <sup>a</sup>	Frequency
<i>Or22a/Or22b</i>	Methyl hexanoate	1	C-	0.053
<i>Or22a/Or22b</i>	Methyl hexanoate	2	TI	0.105
<i>Or22a/Or22b</i>	Methyl hexanoate	3	T-	0.842
<i>Or35a</i>	Methyl hexanoate	1	AA	0.158
<i>Or35a</i>	Methyl hexanoate	2	GG	0.053
<i>Or35a</i>	Methyl hexanoate	3	GA	0.789
<i>Or35a</i>	Ethyl hexanoate	1	AA	0.158
<i>Or35a</i>	Ethyl hexanoate	2	GG	0.053
<i>Or35a</i>	Ethyl hexanoate	3	GA	0.789
<i>Or47a</i>	Ethyl hexanoate	1	TTCAATC	0.132
<i>Or47a</i>	Ethyl hexanoate	2	GTCACDT	0.053
<i>Or47a</i>	Ethyl hexanoate	3	GCTGCDT	0.079
<i>Or47a</i>	Ethyl hexanoate	4	TCTGCDT	0.026
<i>Or47a</i>	Ethyl hexanoate	5	GTCAATC	0.711

<sup>a</sup>An "I" indicates an insertion polymorphism and "D" indicates a deletion polymorphism.

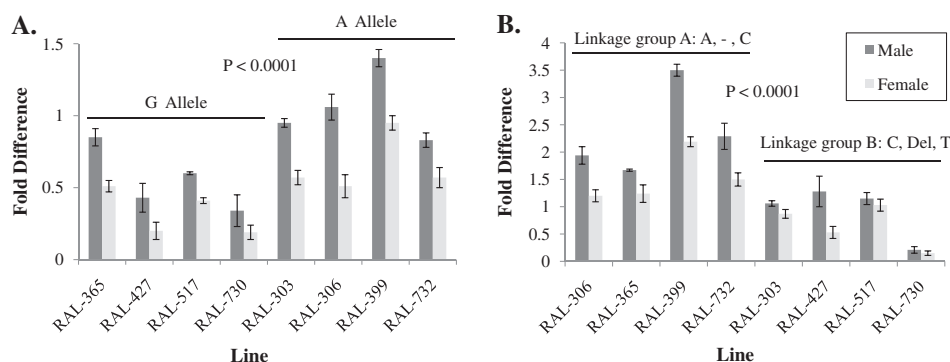
**Figure 6** Haplotype analysis of polymorphisms significantly associated with behavioral responses to (A) methyl hexanoate or (B) ethyl hexanoate for each odorant receptor locus. Significant behavioral differences among haplotypes were determined by ANOVA. Differing haplotypes were subsequently identified by post hoc Tukey's test at  $P < 0.05$  and indicated by unique letters above the bars.

to ethyl hexanoate (Figure 7). Specifically, increased *Or35a* expression was associated with lines carrying the -455A allele that had overall positive behavioral response indices ( $df = 1$ ,  $F = 41.40$ ,  $P < 0.0001$ ). In the case of *Or47a*, an A1336C polymorphism and 2 markers (1349Del2bp and C1357T) are in the noncoding region and are in complete LD. Lines carrying the linkage group consisting of 1336C, 1349 Del 2 bp, and 1357T are associated with more positive behavioral response indices and decreased levels of *Or47a* expression ( $df = 1$ ,  $F = 54.72$ ,  $P < 0.0001$ ). No significant allele by sex interactions was found.

## Discussion

Multiple polymorphisms were identified in the *Or22a/Or22b*, *Or35a*, *Or47a*, and *Or10a* gene regions that were associated with behavioral variation in response to esters in a natural *D. melanogaster* population. We chose to examine these genes from the family of 60 genes that comprise this multi-gene *Or* family based on previous work elucidating their respective electrophysiological response profiles (Hallem et al. 2004; Hallem and Carlson 2006). *Or22a* and *Or35a* are broadly tuned receptors responding to a large suite of odorants including esters, with methyl hexanoate and ethyl hexanoate both eliciting excitatory responses for *Or22a* expressing neurons and inhibitory responses for *Or35a* expressing neurons. *Or10a* exhibits strong excitation to aromatic compounds with polymorphisms in *Or10a* implicated in behavioral variation in responses to the aromatic compounds, benzaldehyde and acetophenone (Hallem et al. 2004; Hallem and Carlson 2006; Rollmann et al. 2010). *Or10a* has also been shown to be inhibited by several odorants, including ethyl hexanoate. *Or47a* is narrowly tuned and responds chiefly to esters. Thus, our results identifying polymorphisms in these receptors, which were associated with variation in behavioral responses to ethyl hexanoate and methyl hexanoate, are consistent with the role of these receptors in the detection of esters. Also consistent with these results is the lack of an association between polymorphisms in *Or43a* and *Or67b* and behavioral responses to these esters as these receptors have not been shown to have ligand affinity to either ester (Hallem and Carlson 2006). Finally, our results are in line with the combinatorial nature of *Ors*: Multiple receptors responding to a single odorant and single receptors responding to a suite of odorants.

Given that a significant correlation was found between independent behavioral measurements of responses to ethyl hexanoate and methyl hexanoate, we predicted that the association profiles for both odorants would be globally similar. Indeed, we observed striking concordance in the association profile patterns for the 2 odorants and variation in the *Or22a/Or22b* region and *Or35a* region (Figure 5A–B). The association of these polymorphisms with independent behavioral measurements of responses to each of these structurally similar odorants serves to further validate their



**Figure 7** *Or35a* and *Or47a* gene expression levels. For each gene, lines with overall more negative behavioral response indices to ethyl hexanoate and carrying a given allele and lines with more positive response indices carrying the alternate allele were selected. **(A)** *Or35a* expression levels across lines carrying either the −455G allele or the −455A SNP. **(B)** *Or47a* expression levels for the linkage group comprised of A1336C, 1349 Del 2 bp, and C1357T that are in complete LD. A dash indicates a lack of deletion.

significant contributions to variation in odor-mediated behavior. In the case of *Or47a*, individual polymorphisms were associated with only one but not the other odorant raising the possibility that *Or47a* may be particularly adept at discriminating among these esters. The *Or47a* electrophysiological response profile is narrower in its tuning compared with *Or22a* and *Or35a* response profiles (Hallem and Carlson 2006). In addition, since the concentrations for optimal resolution of behavioral variation among wild-derived lines differed for ethyl hexanoate and methyl hexanoate, *Or47a* selectivity may be mediated by concentration-dependent changes. It should be noted that we have examined 7 receptors out of the multigene family and that variation in additional receptors may also contribute to differences in behavioral responses to these odorants. The statistical power for detecting genotype–phenotype associations increases as sample size increases. Therefore, increasing sample sizes within a population and/or surveying additional populations could potentially identify additional polymorphisms associated with variation in behavioral responses to ethyl hexanoate and methyl hexanoate.

Both coding and noncoding changes were associated with variation in behavioral responses to both odorants. Protein coding changes can result in changes in protein structure/function, and amino acid substitutions in vertebrate chemosensory receptors have been shown to change ligand-binding affinity and thereby result in differences in odorant sensitivity (Reed et al. 2004; Nie et al. 2005; Abaffy et al. 2007; Keller et al. 2007). Thus, the Asn348Ser substitution in *Or35a* and Gly396Asp substitution in *Or10a* are of particular interest. The substitutions in both genes are predicted to be located extracellularly and thus could influence ligand binding. Finally, it is possible that polymorphisms may affect dimerization with Orco, a receptor that acts to localize odorant receptors in the sensory dendritic membrane (Jones et al. 2005; Benton et al. 2006) and plays a role in the formation of ligand-gated nonselective cation channels (Sato et al. 2008; Wicher et al. 2008).

Noncoding nucleotide changes can also be primary agents of behavioral variation (Wang et al. 2007, 2010; Rollmann et al. 2010). Noncoding odorant receptor polymorphisms have been previously associated with variation in behavioral responses to benzaldehyde and acetophenone, and high behavioral responder haplotypes to benzaldehyde were associated with alterations in odorant receptor gene expression levels (Rollmann et al. 2010). SNPs in noncoding regions of odorant-binding proteins contributing to olfactory behavior also have been implicated in alterations to messenger RNA (mRNA) secondary structure and gene expression levels (Wang et al. 2007, 2010). In *D. melanogaster*, ~500-bp region upstream of *Or* genes has been shown to be sufficient for maintaining appropriate OR expression patterns (Ray et al. 2007). Furthermore, misexpression of odorant receptors has been shown to result in altered olfactory avoidance behavior (Stortkuhl et al. 2005). In our study, the majority of polymorphisms associated with variation in olfactory behavior were noncoding changes. These polymorphisms were not located in promoter motif regions identified to date (Ray et al. 2007, 2008) nor in conserved motifs identified in 3' untranslated gene regions. However, consistent with differences in *Or* gene expression levels contributing to variation in behavior, we observed significant differences in *Or35a* and *Or47a* gene expression levels among lines with more positive versus negative overall behavioral responses to ethyl hexanoate. Of particular note is the G-455A *Or35a* marker associated with variation in behavioral response to both methyl hexanoate and ethyl hexanoate, and in which increased expression was associated with more positive response indices to ethyl hexanoate. This polymorphism is located directly upstream of the initiation codon between 2 highly conserved blocks of sequence, suggesting it could contribute to the regulation of gene expression. Moreover, the *Or47a* polymorphisms (1349Del12bp and C1357T) are located within 12 bp from the termination codon and thereby could affect regulation of mRNA stability. These changes in gene expression can contribute to differences in odorant sensitivity and discrimination,

with the combinatorial group of odorant receptors together influencing variation in olfactory behavior.

Finally, a significant difference in olfactory sensitivity to these esters has been previously observed among *Drosophila* species (Stensmyr et al. 2003; Dekker et al. 2006), but within-species variation had yet to be examined. Our results show that significant variation in sensory perception exists within a natural *D. melanogaster* population for ethyl hexanoate and methyl hexanoate and that variation in the *Or22a/Or22b* gene region as well as in *Or10a*, *Or35a*, and *Or47a* contributes to this variability in behavioral response. Thus, changes at the most peripheral sensory stage can contribute to within-species differences in behavioral responses to environmental stimuli.

## Supplementary material

Supplementary material can be found at <http://www.chemse.oxfordjournals.org/>

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